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U.S. National Phase Patent Application Attorney Docket Number: 18016-23 Express Mail Number: EQ 336 718 587 LIS

MODULATION OF PROTEIN METHYLATION AND PHOSPHOPROTEIN PHOSPHATE

# **CROSS-REFERENCE TO RELATED APPLICATIONS**

[0001] This application claims the benefit of priority to PCT/US03/07658, filed March 13, 2003, which claims priority to U.S. provisional application No. 60/363,537, filed March 13, 2002. The entire contents of each of these applications is incorporated herein by reference in its entirety.

#### **BACKGROUND**

[0001] A variety of diseases are characterized partly by alteration in the pattern or amount of phosphate in regulatory or structural proteins. Protein phosphate content generally is controlled by phosphate addition, which is catalyzed by kinases, and by phosphate removal, which is catalyzed by phosphatases. Whereas kinases most often are regulated with great specificity, protein phosphatases are characteristically less selective. Thus, whereas kinases generally trigger systemic events in response to rather singular, specific stimuli, and then generally do so only through one signaling pathway, phosphatases exhibit broad dephosphorylating activity toward many of the phosphoproteins in their environment. Because of their specificity, kinases have always seemed especially attractive targets for drug development. For exactly the same reason, phosphatases, because of their lack of selectivity and broad systemic activities, have been viewed unfavorably as drug development targets.

[0002] Nevertheless, a number of attempts have been made to use phosphatase modulators for therapeutic purposes. For instance, Schieven in US patents Nos. 5,565,491 and 5,693,627 reports on the use of phosphotyrosine phosphatase inhibitors to control proliferation of immune B cells. The patents disclose inhibitors that act directly on the phosphatase: metal-organic coordinate covalent compounds, nonhydrolyzable phosphotyrosine analogs, the streptomyces protein phosphatase inhibitor Dephostatin, and the prostatic acid phosphatase inhibitor 4-(fluoromethyl)phenyl phosphate. The inhibitors, as disclosed in the patents, inhibited

proliferation of B cell leukemia and lymphoma cells, but also inhibited proliferation of normal B

cells.

[0003] Lazo et. al. in US patent Nos. 5,700,821, 5,856,506, and 5,925,660 discloses synthetic

phosphatase inhibitors produced by combinatorial synthesis using L-glutamic acid as the initial

scaffold. As disclosed in the patents, the compounds inhibited a variety of protein phosphatases,

including PP1, PP2A, PP3, CDC25A, and CDC25B, and inhibited proliferation of human breast

cancer cells in culture.

[0004] Hemmings discloses in US patent No. 6,159,704 modulation of the phosphatase activity

of the catalytic subunit of PP2A ("PP2Ac") via its interaction with eRF1. As disclosed, eRF1 is

the ribosome-associated factor responsible for polypeptide chain release at the termination of

protein synthesis; but, it also binds to and interacts with the catalytic subunit of PP2A.

According to the disclosure, eRF1 recruits PP2A to the ribosome and mediates the role of PP2A

in protein synthesis. According to the patent, the inhibitors disrupt the interaction between eRF1

and PP2Ac. As further disclosed in the patent, they, thus, might inhibit protein synthesis, and

therefore, might be useful to reduce aberrantly high protein synthesis and cell proliferation

which, accordingly, might make them useful for treating proliferative disorders.

[0005] Yet another example in this regard is disclosed by Honkanen et. al. in US patent No.

5,914,242. As disclosed in the patent, inhibitors of certain serine/threonine protein phosphatases,

in particular fostriecin, an organic compound first isolated from streptomyces, is used to reduce

damage to the heart following myocardial infarction. According to the patent, fostriecin inhibits

PP2A thereby causing greater phosphorylation of the protein 1-2. This leads to proteolysis of 1-

2 and reduces its level in the cell, because the phosphorylated protein is a much more active

substrate for the protease. Since 1-2 inhibits PP1, the decrease in 1-2 activity due to proteolysis

results in increased PP1 activity. According to the disclosure, increased PP1 activity protects

cells from the deleterious effects of ischemia, although the mechanism of protection is not

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known. Further according to the disclosure, the protective effect of fostriecin might be due to

inhibition of phosphatase activity that results in less dephosphorylation of proteins

phosphorylated by protein kinase C.

[0006] The inhibitors in all of the foregoing patent disclosures, except Honkanen, act directly on

the phosphatase, inhibiting its activity competitively or irreversibly. The inhibitor disclosed by

Honkanen acts specifically to disrupt the interaction of the phosphatase, PP2Ac, with a

ribosomal protein and likely will affect primarily the action of PP2Ac on protein synthesis, rather

than its more general action as a phosphatase. In any case, all of the inhibitors of the foregoing

patents act solely to decrease the activity of phosphatases. Inherently they cannot act to increase

phosphatase activity, although, this is desirable in many cases.

[0007] Alzheimer's Disease (AD) is a progressive neurodegenerative disease associated

clinically with memory impairment and decreased cognitive function [Selkoe, 2001 #2]. Post-

mortem brains of AD patients display two pathological hallmarks: neuritic plaques and

neurofibrillary tangles (NFTs). The plaques are extracellular deposits. They are composed of

amyloid b-protein (Ab), which is a peptide derived from proteolytic cleavage of the amyloid

precursor protein. NFTs, in contrast, are found primarily within the cell body. They are

composed, in large part, of filaments of tau protein.

[0008] Tau normally is found predominantly in the axons of neurons where it stabilizes

microtubules (MTs) and promotes their polymerization [Buee, 2000 #3]. MTs play a major role

in maintaining the cellular architecture of neurons and are largely responsible for axonal

transport [Goldstein, 2000 #4]. The integrity of MT structure is therefore critical for proper

neuronal function and synaptic transmission. While tau normally is phosphorylated, it is

abnormally hyperphosphorylated in NFTs [Grundke-lqbal, 1986 #47]. Increased

phosphorylation appears to precede and promote NFT formation [Alonso, 2001 #5][Alonso,

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1996 #10]. Hyperphosphorylated tau is also found in the cytobsol of NFT-containing neurons

[Kopke, 1993 #49]

[0009] Phosphorylation inhibits tau's ability to bind and stabilize MTs [Bramblett, 1993

#7][Biernat, 1993 #8][Alonso, 1994 #9]. Furthermore, hyperphosphorylated tau has a dominant

negative effect in that it promotes MT disassembly by binding normal tau, MT associated protein

1, and MT associated protein 2, interfering with the ability of these three proteins to stabilize

MTs [Alonso, 1997 #48]. These effects help account for the observation that neurons containing

NFTs lack MTs. The cytoskeletal disruption brought about by hyperphosphorylated tau thus

provides an explanation for its role in the neurodegeneration associated with AD.

[0010] Genetic evidence supports the conclusion that a critical event in the development of AD-

type dementia is tau hyperphosphorylation [Lee, 2001 #6]. Though under some conditions Ab

accumulation has been shown to promote NFT formation [Lewis, 2001 #11][Gotz, 2001 #12],

plaque formation is not essential for NFT-associated dementias, the so-called 'tauopathies'.

Mutations in the tau gene underlie several familial neurodegenerative diseases where filamentous

deposits of hyperphosphorylated tau have been observed in the absence of amyloid plaques, most

notably fronto-temporal dementia and Parkinsonism linked to chromosome 17 [Lee, 2001 #6].

[0011] Tau hyperphosphorylation results from an imbalance between kinase and phosphatase

activities. Phosphorylation is catalyzed by the neuronally enriched serine/threonine kinases

glycogen synthase kinase 3b (GSK-3b) and cyclin-dependent kinase 5 (CDK5) [Buee, 2000]

#3][Billingsley, 1997 #13]. The most important tau dephosphorylating enzyme is protein

phosphatase 2A (PP2A) [Planel, 2001 #14][Merrick, 1997 #15][Kins, 2001 #16][Gong, 1994

#50].

[0012] Recent results suggest that a decrease in PP2A activity, rather than increased kinase

activities, is crucial for the elevated levels of tau phosphorylation associated with NFT

formation. PP2A expression has been found to be significantly reduced in the hippocampus of

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AD brains relative to control brains [Vogelsberg-Ragaglia, 2001 #17], and expression studies in

mouse brain indicate a general decrease in PP2A expression levels with age [Jiang, 2001 #18].

Treatment of cultured human neurons with the PP2A inhibitor okadaic acid results in tau

hyperphosphorylation, reduced binding of tau to MTs, MT depolymerization, and axonal

degeneration [Merrick, 1997 #15]. Moreover, starved mice display a pattern of tau

hyperphosphorylation similar to that found in AD brains [Planel, 2001 #14], and this

hyperphosphorylation appeared to result from decreased PP2A activity towards tau rather than an

increased kinase activity. In fact, the tau phosphorylating activities of CDK5 and GSK-3b

decreased under these conditions. Thus, reduced PP2A activity towards tau must be part of any

model accounting for NFT formation during the progression of AD.

[0013] PP2A is a multimeric protein complex consisting of a 65 kDa A subunit that acts as a

scaffold for the association of a 36-kDa catalytic C subunit and one of a variety of regulatory B

subunits [Janssens, 2001 #19]. B subunits control the substrate specificity and subcellular

localization of PP2A. Ba, the major regulatory subunit in brain [Kamibayashi, 1994 #20], targets

trimeric PP2A to MTs [Sontag, 1995 #21] and dramatically increases the enzyme's activity

towards the tau protein [Sontag, 1996 #22]. ABaC heterotrimers bind directly to the carboxyl-

terminal MT binding domain of tau [Sontag, 1999 #23]. The highly conserved carboxyl-terminal

sequence of the PP2A C subunit is a focal point for the enzyme's regulation. Reversible methyl

esterification of the C-terminal leucine a-carboxyl group of the PP2A C subunit is a major locus

of control [Tolstykh, 2000 #26] [Wu, 2000 #27] [Yu, 2001 #28] [Wei, 2001 #29].

[0014] PP2A methylation is controlled by a specific S-adenosylmethionine (SAM) dependent

methyltransferase [Lee, 1993 #24] and a specific methylesterase [Lee, 1996 #25]. Methylation

modulates PP2A activity by controlling the association of regulatory B subunits with the

catalytic AC core [Tolstykh, 2000 #26][Wu, 2000 #27][Yu, 2001 #28][Wei, 2001 #29]. The

assembly of ABC heterotrimers proceeds as a multistep process with AC dimer methylation

followed by binding of regulatory B subunits (figure 1). Tolstykh et al. [Tolstykh, 2000 #26]

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demonstrated that methylation of AC dimers from bovine brain dramatically increases their

affinity for Ba regulatory subunits. Given the critical role of Ba in targeting PP2A activity

towards tau, a decrease in PP2A methylation could lead to tau hyperphosphorylation, NFT

formation, and neurodegeneration.

[0015] Decreased PP2A activity can contribute not only to tau hyperphosphorylation, but it can

lead to other clinical indicators. For instance, homocysteine, through SAH hydrolase, is an end

product of SAM-dependent methylation. The hydrolase reaction is reversible, and actually

favors condensation of homocysteine and adenosine to form SAH (S-adenylhomocysteine).

SAH is a potent inhibitor of methylation and accumulation of homocysteine (Hcy) thus generally

is accompanied by increased SAH and, consequently, is associated with decreased methylation

activity. Therefore, high plasma homocysteine levels generally may be indicative of decreased

protein methylation and resultant decreases in methyl-dependent protein activities, such as PP2A

phosphatase.

[0016] Indeed, over the last several years data has emerged in the clinical literature

demonstrating a significant correlation between elevated plasma homocysteine (Hcy) and the

occurrence of AD [Seshadri, 2002 #1][McCaddon, 1998 #30][Clarke, 1998 #31]. Elevated

plasma Hcy has long been established as an independent, graded risk factor for cardiovascular

disease [Clarke, 1991 #32][Boushey, 1995 #33][Welch, 1998 #34]; but, its role in AD has taken

longer to establish. An early study found that patients with pathologically confirmed AD had

significantly elevated plasma Hcy levels relative to a control group [Clarke, 1998 #31]. Hcy

levels in the AD patients remained stable over time even as the disease progressed, suggesting

that the elevation was not a result of neurodegeneration. Furthermore, patients with high plasma

Hcy displayed more rapid neural atrophy over the course of three years than did patients with

lower levels. More recent data from a prospective study provides convincing evidence that a rise

in plasma Hcy precedes the onset of AD and is an independent risk factor for the disease

[Seshadri, 2002 #1]. Baseline plasma Hcy levels were measured in 1092 non-demented patients

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and the occurrence of AD in this group was followed for several years. After adjusting for other

AD risk factors, the authors found that plasma Hcy levels greater than 14 micromolar coincided

with a roughly two-fold increased risk of developing AD. Further, elevated plasma Hcy appears

to be a graded risk factor, with a 40 percent increased risk of developing AD associated with

each 5 micromolar incremental rise. These studies clearly indicate a connection between high

plasma Hcy and AD. While it has been recognized that insight into the mechanism underlying

the association could give important clues for treatment of the disease, recognition of the

association thus far has not provided a better understanding of AD or any other disease.

[0017] Elevated plasma homocysteine has been established as a risk factor not only for AD but

also for heart disease, Type 2 diabetes, obesity, multiple sclerosis, stroke, cancer, rheumatoid

arthritis, vascular disease, and birth defects; as well as various neurological illnesses including,

among others, Parkinson's, depression, schizophrenia, and alcoholism. The relationship between

elevated serum homocysteine and underlying disease etiology has not been elucidated for any of

these diseases. Perhaps because of this, establishing the link has not led to effective therapeutic

modalities, as yet. The general situation for all the diseases in this regard is fairly well illustrated

by the foregoing discussion relating to Alzheimer's Disease. Presently nutritional

supplementation is the only intervention thus far available for altering plasma homocysteine

levels, and thereby, perhaps, reducing risk factors for these diseases. Unfortunately, whatever

the efficacy of nutritional intervention at reducing plasma homocysteine, there is no evidence as

yet that nutritional intervention actually reduces the homocysteine associated risk factor for

disease. Furthermore, nutritionally forced reductions in plasma homocysteine actually may have

deleterious effects.

[0018] Clearly, there is a need for an improved understanding of: the link between homocysteine

levels, disease risk factors and disease itself, its etiology, and the factors that control the

development and progress of the diseases. Even more important and pressing is the need for

better diagnostic tools and for, above all, effective therapies for AD. Unfortunately, AD is

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merely illustrative in this regard. Many other diseases also are poorly understood, hard to

diagnose, and presently lacking effective treatments.

[0019] Heretofore, proteins such as PP2A did not seem promising targets for effective

therapeutics. Typically, they are ubiquitous, abundant and, perhaps worse for drug development,

they are important general regulators of protein phosphokinase or protein phosphatase activities

that affect virtually all phosphoproteins, and they interact with a very wide range of regulatory

proteins. Thus, It appeared likely that targeting them would only lead to general systemic

distress. Furthermore, it seemed likely that modulators of phosphatase activity would suffer

some of the same disadvantages as the inhibitors discussed above. These inhibitors

disadvantageously target a broad spectrum of serine/threonine proteases involved in regulating

and performing vital cell processes and, consequently, broadly affect cellular metabolism and

physiology, often with undesirable or deleterious consequences. Given their similar ubiquity and

regulatory role, the same disadvantages were expected to limit the usefulness and efficacy of

agents that modulate the activities of phosphatases. Thus, there has been and there continues to

be a real need for improved diagnostics, better agents for altering the activities of proteins

important in disease process, and effective methods for treating disorders and diseases such as

AD.

SUMMARY OF THE INVENTION

[0020] The present invention, in part to overcome these problems provides among other things

as follows.

[0021] A method for identifying a compound for altering a protein activity, comprising:

identifying a compound that modulates methylation of a protein phosphatase that affects a

protein activity and determining that the protein activity is altered by modulating with the

compound methylation of the protein phosphatase. In particularly preferred embodiments in this

regard the protein phosphatase is a PP2A protein phosphatase. In another regard in certain of the

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preferred embodiments in the protein activity is the phosphate level of a phosphorylated protein.

In certain highly preferred embodiments in these and other regards the protein activity is

phosphorylation of tau. In especially highly preferred embodiments in these and other regards

tau is hyperphosphorylated, and the compound increases methylation of PP2A and decreases tau

hyperphosphorylation.

[0022] In another aspect, certain preferred embodiments of the invention provide methods for

identifying a composition for altering a protein activity, comprising: identifying a composition

that modulates methylation of a protein phosphatase that affects a protein activity, using the

composition to modulate methylation of the phosphatase, and determining that modulating

phosphatase methylation alters the protein activity. In certain aspects of the invention in this

regard, in certain of the preferred embodiments, the composition is an extract of a natural

product. In further preferred embodiments in this regard the composition is an extract of a

traditional medicine. In another preferred aspect of the invention in these and other regards, the

protein phosphatase is a PP2A protein phosphatase. In a particular aspect of the invention

further in these regards, in certain of the preferred embodiments the protein activity is tau

phosphorylation. In especially highly preferred embodiments in this regard, tau is

hyperphosphorylated, the compound increases methylation of PP2A and decreases tau

hyperphosphorylation.

[0023] The invention further provides in certain aspects and preferred embodiments, compounds

for altering protein activity, wherein the compounds are identified by a method comprising:

identifying a compound that modulates methylation of a protein phosphatase that affects a

protein activity and determining that the protein activity is altered by modulating with the

compound methylation of the protein phosphatase. In particularly preferred embodiments in this

regard the protein phosphatase is a PP2A protein phosphatase. In another regard in certain of the

preferred embodiments in the protein activity is the phosphate level of a phosphorylated protein.

In certain highly preferred embodiments in these and other regards the protein activity is

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phosphorylation of tau. In especially highly preferred embodiments in these and other regards tau is hyperphosphorylated, and the compound increases methylation of PP2A and decreases tau hyperphosphorylation. The invention further provides compositions comprising the compounds. In addition, the invention provides methods for treating cells to alter therein an activity of a protein, comprising administering to the cells by an effective route and/or in an effective amount a compound in accordance with the invention and/or a composition comprising a compound in a accordance with the invention.

[0024] In yet another aspect of the invention, certain of the preferred embodiments provide compounds and/or compositions for altering a protein activity, wherein the compound is identified by a method comprising identifying a compound and/or a composition that modulates methylation of a protein phosphatase that affects a protein activity, using the composition to modulate methylation of the phosphatase, and determining that modulating phosphatase methylation alters the protein activity. In another regard in certain of the preferred embodiments in the protein activity is the phosphate level of a phosphorylated protein. In another preferred aspect of the invention in these and other regards, the protein phosphatase is a PP2A protein phosphatase. In a particular aspect of the invention further in these regards, in certain of the preferred embodiments the protein activity is tau phosphorylation. In especially highly preferred embodiments in this regard, tau is hyperphosphorylated, the compound increases methylation of PP2A and decreases tau hyperphosphorylation. In certain aspects of the invention in these and other regards, in certain of the preferred embodiments, the composition is an extract of a natural product. In further preferred embodiments in this regard the composition is an extract of a traditional medicine.. In these regards, further, in certain aspects the invention provides in certain of the preferred embodiments methods for treating cells to alter therein an activity of a protein, comprising administering to the cells by an effective route and/or in an effective amount a compound in accordance with the invention and/or a composition comprising a compound in a accordance with the invention. In particularly preferred embodiments in this regard, in

accordance with certain aspects of the invention, the compound and/or composition is derived

from a natural product, it affects methylation of PP2A and decreases tau hyperphosphorylation

and it is administered to patients suffering from AD.

**BRIEF DESCRIPTION OF THE FIGURES** 

[0025] Figure 1 is a schematic diagram illustrating processes involved in PP2A methylation and

tau phosphorylation. As illustrated in the figure: (a) PP2A methyltransferase (MTase) (also

referred to as PPMT) binds to and methylates AC dimers (AC and ACmeth); (b) a

methylesterase (MEase) (also referred to as PPME) catalyzes the reverse reaction and removes

methyl groups; (c) methylation of AC dimers dramatically increases their affinity for Ba

subunits, resulting in increased formation of ABaCmeth trimers; (d) the ABaC[meth] timer is the

major tau phosphatase activity in vivo; (e) whereas phosphorylation appears to be controlled

primarily by GSK-3b and CDK5. Increased methylation of PP2A AC dimers leads to increased

amounts of ABaCmeth, increased phosphatase activity and decreased levels of tau

phosphorylation. (SAM: S-adenosylmethionine)

[0026] Figure 2 is a schematic diagram of key steps and enzymes in the methyl cycle. Enzymes

are indicated by circled numbers as follows: (1) methionine adenosyltransferase; (2) SAM-

dependent methyltransferases; (3) SAH hydrolase; (4) cystathionine b-synthase; (5)

cystathionine g lyase; (6) betaine homocysteine methyltransferase; (7) methionine synthase.

DESCRIPTION

[0027] The key regulatory roles of protein kinases and protein phosphatases are well known.

Enzymes of both types have been and continue to be the primary drug development targets.

Although these efforts have met with some success, they have not produced, as yet, many hoped

for advances. The invention herein disclosed in certain of its preferred aspects and embodiments

relates to bioactive agents and methods for modulating the activity of regulatory enzymes, such

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as kinases and phosphatases, without incurring difficulties that have been encountered with other

approaches.

[0028] Bioactive agents that act directly on key kinases and phosphatases, at least at present,

generally act as inhibitors, and their effects do not discriminate among the varied activities of

these enzymes, which generally carry out a variety of regulatory functions. Furthermore, the

effects of the agents cannot be calibrated with precision. Consequently, direct-acting inhibitory

agents too often not only reduce the unwanted activity but also, by the same action, alter the

activity to another aberrant level that also is deleterious.

[0029] Preferred embodiments in some aspects of the present invention augment direct targeting

agents and methods for modulating the activities of enzymes, such as regulatory enzymes,

especially key regulatory enzymes, and overcome some of their disadvantages. In this regard, in

certain aspects of the invention, the preferred embodiments relate to regulatory enzymes that are

kinases or are phosphatases, especially key regulatory kinases and phosphatases, and kinases and

phosphatases that are important determinants of dysfunction and disease.

[0030] In further preferred aspects and embodiments in these and other regards, the invention

relates to indirect targeting for modulating the activities of enzymes, including regulatory

enzymes, especially kinases and phosphatases, particularly kinases and phosphatases that play a

key regulatory role, and those that are important determinants of dysfunction and disease. In

certain highly preferred aspects of the invention in preferred embodiments thereof the invention

provides methods to modulate the activities of target enzymes by acting, partly or entirely, on

other, secondary targets that regulate the activity of the primary target. In certain highly

preferred embodiments of the invention in this regard, the secondary targets naturally regulate

the primary target. Further, in preferred embodiments in this regard, the secondary targets exert

incremental regulatory control on the primary targets. Agents in accordance with this aspect of

the invention thereby are useful to exert incremental control of the activity of the secondary

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target, thus allowing relatively fine control over the primary target activity. In still further preferred aspects of the invention, the preferred embodiments provide both positive and negative

regulatory control of the primary target. In other aspects of the invention in this regard, the

preferred embodiments provide positive regulatory control (stimulatory rather than inhibitory).

Further preferred embodiments provide various combinations of these aspects and embodiments

of the invention.

[0031] In these regards and others, the preferred embodiments in certain preferred aspects of the

invention relate to the activities of primary target enzymes regulated by post-translational

modifications and to modulating the activities of enzymes that effectuate the post-translational

modifications and thereby regulate the activity of the primary target(s).

[0032] These and other aspects of the invention are illustrated by particularly preferred

embodiments relating to the phosphatase PP2A, which plays a central regulatory role in a variety

of cell processes, and to PP2A methyltransferases and methylesterases which, respectively,

methylate and demethylate a specific site in a PP2A subunit. Methylation increases formation of

the active PP2A trimer, thus promoting PP2A phosphatase activity. Demethylation leads the

system back toward the other side of the equilibrium, reducing the amount of active PP2A, thus

decreasing PP2A phosphatase activity. Illustrative of other preferred embodiments of the

invention in this regard, the present invention provides for both up regulating and down

regulating PP2A phosphatase activity, through agents that act, respectively, on PP2A

methyltransferase and PP2A methylesterase.

[0033] The same approach can be applied to modulate the activities of other enzymes, including

among others, kinases and phosphatases. Other activities regulated by methylation also can be

controlled through methyltransferase and methylesterase activities, much the same as described

for PP2A above. Still other activities, not regulated by methylation, but subject to regulatory

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control by other post-translational modifications can be modulated in much the same way with

the other modifications as for the methylation control enzymes discussed above.

[0034] The invention thus relates both to methods for controlling primary target activities,

preferably of primary target enzymes, by effecting control over the activity of a secondary target

activity that controls the activity level of the primary target. It relates further in this regard to

agents for effecting the control, to methods of or for identifying candidate agents, to methods for

verifying the candidates, and for optimizing the agents. It relates as well in these regards and

others to compositions and methods for using the agents to regulate activities in vitro, in cells in

culture, in physiological models, in model organisms, and in treating disorders and disease.

[0035] Furthermore, the invention relates not only to modulating the activity of a primary target

in accordance with the foregoing, but also to the effect of regulation on substrates of the primary

target. An example discussed elsewhere herein in greater detail is illustrative in this regard. In

this example, an agent that inhibits PP2A methylesterase is administered. PP2A methylation

affects primarily Ba-containing PP2A trimers, which are located primarily in brain and liver.

Therefore, the agent will act to increase PP2A phosphatase activity in brain, inter a/ia. In

Alzheimer's Disease decreased PP2A phosphatase activity is associated with marked increase in

tau phosphorylation, and hyperphosphorylated tau is an important component in the

neurodegenerative processes that underlie the disease. The agent, by decreasing demethylation

thereby increasing PP2A phosphatase activity, concomitantly decreases

hyperphosphorylation and ameliorates disease symptoms. The same end result in accordance

with the invention can be accomplished with a variety of other systems involving other primary

target enzymes, other secondary enzymes, and different post-translational modifications.

[0036] In accordance with these and other aspects of the invention, the secondary targets,

exemplified above by PP2A methyltransferase and PP2A methylesterase, provide novel targets

for high throughput screens for agents that can be used not only to regulate the activities of the

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secondary targets themselves, but also thereby the activity of primary targets. Agents that act on

the secondary targets thereby effect control and/or to modulate the activity or occurrence of a

disorder and/or disease associated and/or causative factor. The invention in this regard further

relates in certain of its preferred aspects and embodiments to formulations of such agents for

diagnosis and treatment uses. Further, it will be noted that in accordance with further preferred

embodiments of the invention, PPMS (PP2A methylation status) defects are detected that are

critical early steps in the development of a disorder and/or disease and that, in especially

preferred embodiments in this regard, occur well before the onset of most other symptoms. The

opportunity for very early intervention, thus provided by the invention, enhances agents and

therapies that target the PP2A methylating and demethylating enzymes in accordance with other

aspects of the invention herein discussed and provides thereby the best avenue to effective

preventative and therapeutic protocols for diseases that currently lack effective treatments.

[0037] In certain particularly highly preferred aspects of the invention in this and other regards,

certain preferred embodiments of the invention relate to PP2A, to PP2A comprising methyl-

accepting subunits, to the activities and physiological effector functions of PP2A and to the

modulation of one or more PP2A physiological activities to maintain health, to prevent disorders

and/or disease, to treat disorders and/or disease, and to ameliorate and/or eliminate adverse

symptoms of disorder and/or disease. In particular, the invention relates to PP2A phosphatase

activity and its effect on specific phosphoproteins, especially phosphoproteins regulated by

phosphorylation and/or by dephosphorylation, most particularly those that play an important role

in maintaining health and/or in the development of disorders and/or disease.

[0038] In a particular aspect in this regard, the invention relates to the modulation of PP2A

activity, to the modulation thereby of PPMS, to the modulation of phosphoprotein phosphatase

activity thereby, and to the methyltransferase and methylesterase enzymes that control PPMS,

including, among others, agents that facilitate methylation of PP2A or that inhibit demethylation

of methylated PP2A. In addition, the invention relates to the identification of factors that

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ameliorate defects in PPMS, particularly those that mimic the effects on PP2A phosphatase

activity of PP2A methylation, especially those that activate PP2A phosphatase activity,

particularly PP2A phosphatase activity towards hyperphosphorylated proteins that, when

hyperphosphorylated, cause aspects of disorder and/or disease, including, in particular, the

microtubule-associated protein, tau, which, when hyperphosphorylated is a causative agent of

neurodegeneration and mental illness, among others, often associated with AD.

[0039] The invention further relates in this and other regards to formulations for administering

the aforementioned agents to subjects to increase methylation and activity of PP2A such as

formulations that increase methylation and PP2A dephosphorylation activity toward

hyperphosphorylated proteins, particularly proteins that exhibit hyperphosphorylation in disorder

or diseased physiological states but not in healthy states, such as, in one very highly preferred

embodiment in this regard, hyperphosphorylated tau protein associated with onset of Alzheimer's

Disease and/or with the disease itself.

[0040] In these and other regards, the invention provides agents, formulations, and treatment

strategies for the amelioration of PPMS deficiencies so that individuals can arrest, retard, or

reverse the development of the aforementioned diseases. In this regard, in certain particular

aspects and embodiments, the invention provides agents, formulations, and treatments for

familial defects in metabolism or metabolic regulation that cause disease by increasing protein

phosphorylation or preventing protein dephosphorylation by providing methods to modulate

PP2A phosphoprotein phosphatase activity toward critical phosphoprotein substrates that cause

disease, such as hyperphosphorylated tau protein.

Indicators, metrics, and diagnostics

[0041] Furthermore in accordance with the invention herein disclosed, aspects of methylation

metabolism and PP2A methylation provide quantitative and qualitative diagnostic markers of

disease risk, early indications of disease development, and the presence of disease. The

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invention relates in this regard to, among other things, methods for assessing, measuring,

determining, assaying, and the like, PP2A phosphoprotein phosphatase activity and PPMS,

particularly in clinical samples, and to methods of diagnosis and prognosis based on these

assessments, measurements, determinations, assays, results, etc.

[0042] Accordingly, the invention provides methods to assess PPMS and determine PPMS

predictive or indicative of disorder or disease, particularly states or conditions that lead to the

aforementioned diseases, especially those that are life threatening. Homocysteine level is one

such metric that provides a useful parameter for risk assessment in this regard, not only of

previously known diseases such as heart disease and AD, but also other diseases that involve

characteristic alterations of PP2A methylation. In addition, the invention further provides

measures of methylation metabolism that are more direct than plasma homocysteine levels and

provide more reliable and more cost effective determinations of PP2A methylation status

(PPMS) and PP2A activity than serum homocysteine measurements. The independent measures

of methylation metabolism and PPMS provided by the present invention overcome

disadvantages of the standard techniques for homocysteine serum determination.

[0043] Notably in this regard, for example, homocysteine is central to the methyl cycle (figure 2)

and plays an important role in cellular methylation [Selhub, 1999 #35][Scott, 1998

#36][Finkelstein, 1998 #37]. SAM-dependent methylation reactions result in the production of

S-adenosylhomocysteine (SAH) (reaction 2), which is subsequently cleaved by the enzyme SAH

hydrolase to adenosine and homocysteine (reaction 3). The SAH hydrolase reaction is reversible

with the equilibrium actually favoring the condensation of homocysteine and adenosine to form

SAH. SAH is a potent competitive inhibitor of virtually all methyltransferase enzymes and

accumulation of Hcy is associated, via an increase in SAH, with a global decrease in cellular

methylation [Wang, 1997 #39][Yi, 2000 #40][Caudill, 2001 #41]. These results are consistent

with the hypothesis that Hcy facilitates the progression of AD by inhibiting brain PP2A

methylation. Decreased C subunit methylation will result in reduced ABaC heterotrimer

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formation leading to tau hyperphosphorylation, NFT formation, neurodegeneration, and

dementia. Thus, PP2A methylation links elevated plasma Hcy and AD, and it therefore provides

a more direct and more accurate assay of AD indicators in this regard than serum Hcy. The

invention in this regard provides agents, formulations, and treatments for assessing and reducing

the risks of the aforementioned diseases predicted by measurements of elevated plasma

homocysteine (> 10 micromolar).

[0044] The invention also provides in some aspects and embodiments novel methods for PPMS

assessment that provide the first direct assays to assess the efficacy of agents, formulations, and

treatments designed to ameliorate abnormalities, promote health, and prevent or retard the

development of the aforementioned disease states. Preferred methods of the invention in this

regard include measuring DNA hypomethylation in blood, measuring levels of S-

adenosylmethionine and S-adenosylhomocysteine in blood and/or cerebrospinal fluid, measuring

levels of adenosine, methionine, folate, and homocysteine in blood and/or cerebrospinal fluid,

and measuring levels of protein N-methylation in hair or skin. In accordance with further aspects

of the invention in this regard, PPMS defects are detected that are critical early steps in the

development of these diseases, occurring well before the onset of most symptoms, thus allowing

earlier diagnosis and interventions that are both more timely and, being more timely, also are

more effective.

The determination of PP2A methylation and demethylation

[0045] The invention provides several methods for detecting PPMS, directly, and for

characterizing normal and abnormal PPMS in an organism, particularly in a human. A preferred

method in this regard measures levels of PP2A methyl esterification in whole blood by

immunoassay using antibody specific for methylated PP2A that does not significantly cross-react

with unmethylated PP2A. Antibodies specific for methylated PP2A and other reagents and

methods for use in immunoassays in this regard are described in Tolstykh et a/. (2000);

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"Carboxyl methylation regulates phosphoprotein phosphatase 2A by controlling the association

of regulatory B subunits;" EMBO J. 19: 5682-5691 [#26], which is herein incorporated by

reference in its entirety in parts pertinent particularly to antibodies, reagents, and methods for

determining the presence and extent of methylation of PP2A.

[0046] In addition to the aforementioned immunological assays, PP2A methylation and

methylesterase activity can be determined by methanol release. PP2A can be incubated in mildly

alkaline solution, which hydrolyzes the PP2A methyl ester, releasing methanol. After incubation

to complete hydrolysis, methanol is measured by conventional techniques. Methylesterase-

catalyzed PP2A demethylation is measured much the same way, but the methanol determined is

that released by the methylesterase at neutral pH, not by alkaline hydrolysis. The released

methanol at neutral pH is measured by the same standard procedures.

[0047] The results of PP2A methylation determinations in accordance with the foregoing

methods, or other methods, provide an accurate assessment of PP2A activity, are diagnostic of

PP2A insufficiency and, in some cases, of disease, and are used in accordance with the invention

to diagnose disease, particularly at early stages, to assess the utility and likely effectiveness of

interventions that affect PP2A methylation and activity, and to follow and assess the effects and

efficacy of any such intervention undertaken, and of other treatment modalities as well. The

information from these assays, alone or combined, provides, in accordance with certain

particularly preferred embodiments of the invention in this regard, a clear view of PPMS and of

genetic, behavioral, and/or nutritional components of a PPMS problem. Moreover, in

accordance with these aspects and preferred embodiments of the invention, the results accurately

indicate appropriate remedial treatment modalities and provide a monitor of efficacy throughout

a course of treatment.

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Identification of PPMS-modulating agents

[0048] In another aspect, in certain of its preferred embodiments, the invention further provides

methods to identify and characterize agents to modulate PPMS. In certain preferred

embodiments such agents are identified by screening methods using monoclonal antibodies that

bind specifically to methylated, but not unmethylated PP2A.

[0049] Candidate modulators of PPMS can be screened using the PP2A methylation assays

described above and in the examples. Purification procedures for PP2A methyltransferase and

methylesterase, and additional assays, suitable for use in the invention in this regard, are

described in Lee et al. (1996); A specific protein carboxyl methylesterase that demethylates

phosphoprotein phosphatase 2A in bovine brain; Proc. Natl. Acad. Sci. U.S.A. 93: 6043-6047

[#25] and Lee et al. (1993); Protein phosphatase 2A catalytic subunit is methylesterified at its

carboxyl terminus by a novel methyltransferase; J. Biol. Chem. 268: 19192-19195 [#24], which

both are incorporated herein by reference in their entireties in parts pertinent particularly to the

isolation, purification, assay, and use of the PP2A carboxyl methyltransferase and assay of PP2A

methylation, in particular methylation by PP2A carboxyl methyltransferase.

[0050] As an initial screen, in accordance with certain preferred embodiments of the invention.

candidate agents are added to mixtures containing PP2A, PP2A methyltransferase, and S-

adenosylmethionine methyl donor. Following incubation, the level of PP2A methylation is

assessed using the aforementioned antibody. Alternatively, in addition, methylesterified PP2A is

mixed with the methylesterase and after a period of time the disappearance of methyl groups is

assessed by the disappearance of the methylester epitope measured using the aforementioned

antibody. In a further alternative approach, candidate agents are added to intact cells that express

PP2A, the methylating enzyme, and the demethylating enzyme. Following a period of

incubation, PP2A methylation is assessed. The impact of such treatments on PPMS is then

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directly assessed by measuring PP2A methylation by Western Blot analysis using the

aforementioned antibodies.

[0051] In addition to the aforementioned immunological assays, methanol can be determined to

measure PP2A methylation and demethylation, as described in examples set forth below. PP2A

methylation is determined by incubating PP2A in mildly alkaline solution, which hydrolyzes the

PP2A methyl ester, releasing methanol. After incubation to complete hydrolysis, methanol is

measured by conventional techniques. Methylesterase-catalyzed PP2A demethylation is

measured much the same way, but the methanol determined is that released by the

methylesterase at neutral pH, not by alkaline hydrolysis. The released methanol at neutral pH is

measured by the same standard procedures.

Genetic markers

[0052] In a further aspect, the invention provides in certain preferred embodiments, methods for

genetic analysis of aberrations that cause disease susceptibility, indicate likelihood of developing

a deleterious hereditary condition of PPMS, and/or are predictive of treatment tolerance, side

effects, and efficacy. Certain preferred embodiments in this regard relate to SNP(s) associated

with alterations of PPMS, and to analysis of SNPs by standard techniques, such as specific probe

PCR, LCR, RFLP, PCR-RFLP, and a variety of other methods useful both to discover and to

determine SNPs. Also particularly preferred, in the certain aspects and embodiments of the

invention in this regard, are DNA array analyses, including analyses to identify SNPs useful for

diagnosis and prognosis in accordance with foregoing aspects of the invention, inter alia,

particularly SNPs in genes that encode the following: protein kinases; more particularly,

phosphoprotein phosphatases, most especially in this regard PP2A; enzymes that directly or

indirectly modulate the activities of protein kinases; and, more particularly, enzymes that directly

or indirectly modulate the activities of phosphoprotein phosphatases, particularly in this regard,

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methyltransferases and methylesterases, most especially in this regard PP2A methyltransferases and PP2A methylesterases.

#### **EXAMPLES**

Methylation assay in partially purified extract using 3H incorporation Example 1:

[0053] A cell-free extract is prepared from fresh mammalian brain tissue by homogenizing the tissue in Tris buffer, pH 7.5. The extract is clarified by centrifugation at 10,000 x g for 30 min. Solid ammonium sulfate is added to the supernatant to 30% saturation and a clarifying precipitate is allowed to form for an hour. The solution then is cleared by centrifugation for 30 min at 10,000 x g. The clarified supernatant is recovered and ammonium sulfate is added to 70% saturation. The resulting suspension is incubated overnight. Then the precipitate is collected by centrifugation at 10,000 x g for 30 min. The resulting pellet is dissolved in Tris buffer, pH 7.5, and then further purified by DEAE ion exchange chromatography with stepwise elution. PP2A and the PP2A methylating and demethylating enzymes are eluted from the column in loading buffer containing 0.3 M NaCl.

[0054] PP2A methylating activity is assayed by 3H incorporation using S-Adenosyl[3Hmethyl]methionine as substrate. An aliquot of the eluate is mixed with S-Adenosyl[3Hmethyl|methionine and incubated under conditions favorable to methyltransferase activity. 3Hmethyl-PP2A formed during the incubation is assayed by standard methanol procedures.

Screening for agents that alter PP2A methylation using partially Example 2: purified extracts and H incorporation assays

[0055] Extracts are prepared and purified and assays of PP2A methylation are carried out as described in the preceding example, except that compounds to be tested for effects on PP2A methylation also are added to the reactions. Effects of the compounds are scored relative to control assays.

#### Example 3: Cell-based PP2A methylation assay

[0056] A cell tissue culture assay is also be employed. This screen is for drugs that cause increases in PP2A methylation in SY-SH5Y cells in the absence and/or presence of homocysteine. SY-SH5Y cells are grown to confluence in 36-well tissue culture plates. After the cells have reached confluence, the growth medium is supplemented with extracts prepared from medicinal herbs in the presence and absence of 200 pM homocysteine. After a suitable time (- 3 h), cell proteins are extracted into 2% SDS, diluted, blotted onto PVDF or nitrocellulose membranes, and probed with monoclonal antibodies that recognize specific epitopes for either total or methylated PP2A. Extracts that increase methylation are identified as drug candidates for further study.

#### Example 4: Diets to induce high plasma homocysteine in mice

[0057] Male C57BL/J6 mice were obtained from the Jackson Laboratories (Bar Harbor, Maine) at 4 weeks of age. Mice were fed on a vitamin sufficient or a vitamin deficient diet for 9 weeks. All of the vitamin-sufficient diets contained folate, vitamin B6, and vitamin B12. All of the vitamin deficient diets lacked these vitamins. One group of mice was fed a vitamin deficient diet that had an enriched methionine content (Harlan Teklad TD97345). Another group was fed a vitamin deficient diet that contained basal levels of methionine (Harlan Teklad TD00428). The same was true for mice on the vitamin-sufficient diet. One group was fed a diet enriched for methionine (Harlan Teklad TD98002) and another was fed standard rodent chow 5001 c (Lab Diets). The mice were allowed free access to both food and water.

# Example 5: Effects of vitamin deficiency on body, heart and brain weight of C57BUJ6 mice

[0058] Wild type mice were placed on diets containing or lacking folate, vitamin B6, and vitamin B12 as described above for nine weeks. Brain and heart weights of mice in each group then were determined as follows. The brain was removed from each animal and frozen in liquid nitrogen prior to recording the weight. The heart was removed from each animal, perfused with

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approximately 10 ml of ice cold phosphate buffed saline, drained of buffer, and frozen in liquid nitrogen prior to recording the weight. Mice fed vitamin-deficient diets had dramatically decreased body and heart weights relative to mice fed vitamin supplemented diets, but not brain weight, as shown below.

Body, brain, and heart weights for mice fed diets with or without folate, vitamin B6, and vitamin B12

	With Folate, B <sub>6</sub> B <sub>12</sub>	Without Folate, B <sub>6</sub>	With/Without
		B <sub>12</sub>	
Body (g)	$25.9 \pm 0.3$	$13.9 \pm 0.4$	1.87
Heart (mg)	115 ± 3	$63 \pm 2$	1.83
Brain (mg)	$397 \pm 8$	$388 \pm 5$	1.02

[0059] Weights are reported ± standard error. N equals 30 for the diets with vitamins and 27 for the diets without vitamins.

[0060] Although the body and heart weights were dramatically reduced by the vitamin deficiencies, the brains appeared to be unaffected. This fits the idea that the brain is a privileged organ with a high priority for residual levels of folate and the B vitamins. One possible explanation is that the brain is somehow a high-priority organ and receives whatever precious folate and В vitamin resources present. Although mice deficient are methylenetetrahydrofolate reductase have decreased amounts of 5-methyltetrahydrofolate in both brain and liver, the brain/liver ratio is over three-fold higher in null mutant mice [Chen, 2001 #51]. These data suggest that folate levels in the brain are maintained at the expense of other organs

[0061] These results are similar to those obtained in previous research relating to the effects of vitamin deficiency on mice deprived of folate [Gospe, 1995 #52] and vitamin B6 [Bender, 1990 #57; Ha, 1984 #58]. Somewhat different results obtained in one study also are worth noting,

formation in ApoE null mice (backcrossed ten generations into C57BUJ6 mice). The mice were

four weeks old in that study, as in the present case, and they were fed the same four diets as used

in this example, but for eight weeks rather then nine. However, that study found that the body

weight of mice fed diets that contained folate, vitamin B6, and vitamin B12 after the eight weeks

did not differ significantly from the body weight of mice fed on diets that lacked these vitamins.

Example 6: Determination of SAM and SAH,

[0062] SAM and SAH levels were measured by high performance liquid chromatography

(HPLC) according to the method described by Fu et al. in "Interrelations between plasma

homocysteine and intracellular S- adenosylhomocysteine;" Biochem Biophys Res Commun 271:

47-53 (2000), which is incorporated herein by reference in its entirety particularly in parts

pertinent to HPLC assays of SAM and SAH.

[0063] Samples of mouse brain were weighed and homogenized in two times weight/volume of

0.5 HCIO4. Samples were centrifuged at 12,500 g for 5 min and the supernatant was filtered

through a 0.22 mm membrane. Aliquots of 25 ml were injected and run on a hypersil 5 mm

particle size, c18 (ODS), 250 x 4.6 mm (Phenomenex) column with an isocratic elution at 1.3

ml/min. The mobile phase consisted of 0.10 mM sodium acetate, 2.4 mM heptanesulfonic acid,

4.2% acetonitrile, and 50 mM sodium perchlorate. The pH was adjusted to 3.5 with 70%

perchloric acid.

[0064] The retention times for SAM and SAH are 23 min and 29 min, respectively. Standards

were run at five different concentrations: 10,000, 5,000, 2,000, 1,000, and 500 pM for SAM, and

10,000, 5,000, 1,000, 100, and 50 pM for SAH. Linear curves, obtained from the SAM and SAH

standards, were used to calculate concentrations of SAM and SAH in the brain homogenates.

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#### Example 7: Determination of plasma homocysteine

[0065] Homocysteine was measured by HPLC in serum samples reduced with tri-n-butylphosphine.

[0066] Serum was prepared from whole blood, which was allowed to clot for 30 min on ice and then centrifuged. Samples were aliquoted, snap frozen and stored at -80°C for analysis. For analysis, 150 ml of serum was thawed and reduced in a mixture containing 50 ml of 0.1 M potassium borate, 2mM EDTA, pH 9.5, 0.2mM mercaptopropionylglycine, and 20ml of 100ml/I of tri-n-butylphosphine in dimethylformamide. The reduction was allowed to proceed for 30 min on ice after which samples were mixed with 125 ml of 0.6 M perchloric acid containing 1 mM EDTA and allowed to react at room temperature for 10 min, and then centrifuged at 15500 g for 10 min. 100 ml was withdrawn from the middle of the supernatant and mixed with 200 ml of 0.2M potassium borate, pH 10.5, containing 5mM EDTA. 100 ml of 1.0 gm Ammonium 7fluorobenzo-2-oxa-1,3-diazole-4-sulphonate/I.OL 0.2M potassium borate, pH 9.5, was added and the mixture incubated at 60°C for 60min. Samples were cooled and then analyzed using HPLC. 10 ml of derivatized sample was used for each injection. Buffer A (0.1 M acetate buffer, pH 4.0, containing 20m1/I methanol) and Buffer B (0.1 M acetate buffer, pH 6.0, containing 50ml/I methanol) were run at a flow rate of 1.0 ml/min from A to B over 12.5 min then 2.5 min of B, a 3.0 min gradient back to A and 2.0 min A before next injection. A standard was also analyzed to quantify values of homocysteine. Mercaptopropionylglycine was added to the reduction buffer as an internal standard to monitor any potential loss during reduction.

## Example 8: Effects of diet on SAM, SAH, Hcy and methylation rate

[0067] Mice were fed on diets containing or deficient in folate, vitamin B6 and B12 as set out above. SAM, SAH and Hcy were determined as described above.

[0068] Brain SAM levels were slightly higher in mice fed on the vitamin deficient diets than mice fed the normal diets. Plasma Hcy and brain SAH were substantially increased inmice fed

the vitamin-deficient diets, compared to mice on the normal diet. The increases in SAH cause the methylation index (SAM/SAH) [Cantoni,1978 #53] to be severely depressed, which reduces the overall methylation rate. The reduction in methylation rate caused by elevated SAH, may be the reason that SAM levels are elevated under nutrient conditions where SAM synthesis would be expected to be reduced.

Status of methylation metabolism in mice fed diets with or without folate, vitamin B6, and vitamin 812

	Folat	Folate: B <sub>6</sub> , B <sub>12</sub>	
	With	Without	
Hcy (pM)	$20.8 \pm 2.7$	$371 \pm 64$	
SAM (nmol/g)	$13.3 \pm 1.0$	$17.0 \pm 1.3$	
SAH (nmol/g)	$1.7 \pm 0.4$	$16.7 \pm 4.4$	
SAM/SAH	$10.0 \pm 1.8$	$1.8 \pm 0.8$	

[0069] Values are reported ± standard error. N=7 for the Hcy value given for mice fed a vitamin-deficient diet, while N=6 for all other values.

Example 9: Determination of PP2A methylation

[0070] SH-SY5Y cells were propagated in a 1:1 mixture of Eagle's Minimum Essential Medium and Ham's F12 medium supplemented with 10% Fetal Bovine Serum, Penicillin (100 units/ml) /Streptomycin (100 mg/ml) and 2 mM L-Glutamine. The flasks were incubated at 37°C (5% CO2). Media was changed every 4-6 days until the cells were confluent. On confluency, cells were treated with 25-200 mM homocysteine for 3 hours. After 3 hrs, the cells were harvested using cold PBS, centrifuged, resuspended in SDS sample buffer, and loaded on a 12.5% percent SDS-PAGE gel. Gels were run at 200 V, transferred onto PVDF membrane at 100 V for 1 hr. Each membrane was blocked for 1 hr in 5% milk, followed by an hour incubation in antibodies, 6A3 (1:50) and 4D9 (1:25) which recognize total and methylated PP2A, respectively. The membranes were then washed 5x for 5 mins each with TBS containing tween (TBST) and then incubated with goat anti-mouse IgG horseradish peroxidase coupled secondary antibody, diluted

1:25,000 in 5% milk prepared in TBS, for 1 hr. Following the secondary antibody incubation, the membranes were washed 5x for 10 mins in TBST and developed with ECL plus (Amersham). Signal intensities of X-ray film exposures for each membrane were quantified by scanning densitometry using NIH Image.

## Example 10: Inhibition of PP2A methylation by homocysteine

[0071] Inhibition of PP2A methylation by homocysteine was determined in cultured cells. Total PP2A and methylated PP2A was determined by western blotting in SH-SY5Y neuroblastoma cells. Homocysteine was added to confluent SH-SY5Y cells to concentrations of 0, 25, 50, 100, and 200 pm. Three hours after adding homocysteine the cells were harvested into cold PBS, mixed with SDS sample buffer, and then analyzed by SDS-PAGE and western blotting. To determine total PP2A, the blots were probed with monoclonal antibody 6A3, which recognizes total PP2A. To determine methylated PP2A the blots were probed with monoclonal antibody 4D9, which recognizes methylated but not unmethylated PP2A. Total PP2A remained much the same at all levels of homocysteine, whereas the level of methylated PP2A declined dramatically with increasing homocysteine concentration.

#### Example 11: Determination of Tau phosphorylation

[0072] Mice brains were homogenized in 10 times volume/weight buffer containing:

[0073] 62.5 mM Tris-HCI, pH 6.8, 10% (w/v) glycerol, 5% (v/v) [3-mercaptoethanol, 2.3% (w/v) SDS, 100 nM okadaic acid (OA), 1 mM phenylmethylsulfonyl fluoride, and 1 mM EDTA. This is a modified version of the buffer reported by Planel et al. [Planel, 2001 #54]. The homogenates were boiled for 5 mins and then centrifuged for 15 mins at 4°C in a Fisher microcentrifuge model 235A. The supernatant was then diluted 1:2 in SDS-sample buffer and loaded on a 12.5% percent SDS-PAGE gel. Gels were run at 200 V, transferred onto PVDF membrane at 100 V for 1 hr. Each membrane was blocked for 1 hr in 5% milk, followed by overnight incubation in antibodies, CP13 (1:200) and PHF1 (1:500) which recognize phosphoserine 202

and phosphoserine 396/404 respectively [Greenberg, 1992 #55; Otvos,1994 #59] and TG5 (1:1000), which recognizes a phosphorylation independent epitope on tau [Jicha,1997 #56]. The following day, membranes were washed 5x for 5 mins each with TBS containing tween (TBST) and then incubated with goat anti-mouse IgG horseradish peroxidase coupled secondary antibody, diluted 1:25,000 in 5% milk prepared in TBS, for 1 hr. Following the secondary antibody incubation, the membranes were washed 5 x for 10 mins in TBST and developed with ECL plus (Amersham). Signal intensities of X-ray film exposures for each membrane were

quantified by scanning densitometry using NIH Image.

Example 12: Effects of vitamin deficiency on tau phosphorylation

[0074] Phospho-tau is primarily dephosphorylated by a heterotrimeric phosphoprotein

phosphatase 2a variant, ABaC. The assembly of ABaC depends on carboxyl methylation of the

catalytic subunit (C). Lowered levels of PP2A methylation thus should lead to lowered rates of

phospho-tau dephosphorylation. Moreover, the protein kinases that are responsible for tau

phosphorylation are themselves activated by phosphorylation. Heterotrimeric, methylated forms

ofPP2A catalyze the dephosphorylation and inactivation of these kinases. Under conditions

where PP2A is disabled because of a methylation deficiency, the kinases would tend to be

activated, which would further contribute to tau hyperphosphorylation. This is demonstrated

using the mouse diet model described below.

[0075] Mice were fed on diets containing or deficient in folate, vitamin B6, and vitamin B12 as

described above. Tau phosphorylation was determined in the mice by SDS-PAGE and western

blotting using the procedures described above. The western blots were probed using three

different tau-specific monoclonal antibodies, as discussed above. As shown in the data below,

the total amount of tau, determined by Western blots probed with TG5 antibody, was essentially

the same for all the diets. Tau phosphorylation, however, was significantly greater in mice fed

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diets deficient in folate, B12, and B6 then in mice fed diets containing these vitamins in normal

amounts.

Tau hyperphosphorylation in mice on diets deficient in folate, B12, and 86

[0076] Date from representative Western blots of extracts prepared from mice raised on normal

diets (A and B) and vitamin-deficient diets and D). CP13 and PHF1 are monoclonal antibodies

specific for phosphorylated tau epitopes. TG5 is a monoclonal specific for Tau independent of

its phosphorylation state. Blots were treated as self-contained data sets.

[0077] Relative Tau (Vitamin deficient / Antibody Normal diet)

[0078] Relative Tau - the fold-induction in the signal relative to the mouse fed standard rodent

chow (A diet) - was calculated for each blot, using the program NIH image. Each value is the

average of 8 determinations, and the  $\pm$  value is the standard error of the average.

Example 13: Effect of methionine enrichment on tau phosphorylation

[0079] The effects of methionine enrichment were determined either in the presence or absence

of a deficiency for folate, vitamin B6, and vitamin B12. When excess methionine was added to

the normal diet, an approximately two-fold increase in plasma Hcy was observed while other

variables showed little difference. In contrast, excess methionine added to vitamin-deficient

diets did not have a sizeable significant effect on tau hyperphosphorylation and methylation

metabolism.

[0080] Effects of methionine enrichment on mice fed diets with or without folate, vitamin B6,

and vitamin B12

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[0081] Ratios are given for values obtained from mice fed diets enriched with methionine divided by values obtained from mice fed diets not enriched for methionine.

	Folate. B <sub>6</sub> , B <sub>12</sub>		
	With	Without	
Нсу	1.9	0.9	
SAM	0.9	1.2	
SAH	0.8	1.1	
CP13	1.1	1	
PHF1	1.2	0.9	
TG5	1.2	1	
Body	0.9	0.9	
Brain	1	1	
Heart	0.9	0.8	

[0082] Feeding and assays were performed as described above. Hey, total plasma homocysteine; SAM and SAH values measured in brain homogenates. CP13 and PHF1, tau phosphorylation and TG5, total tau (see data below) body, brain, and heart weights (see data in examples above).

Example 14: Increased PP2A methylation prevents or ameliorates vascular inflamation and CAD development in hypercysteinimic mice

[0083] In one example, mice are placed on diets that results in hypercysteinemia, and allowed to become hypercysteinemic, as described by Hofmann et al. (March 2001), Hypercysteinemia enhances vascular inflamation and accelerates atherosclerosis in a murine model, J. Clin. Invest. 107(6): 675-683, which is incorporated herein by reference in parts pertinent to hypercysteinemia in the murine model and its use in assay and/or discovery and/or development of agents and/or methods and/or treatment regimens and the like to prevent, treat, ameliorate, retard, reverse or cure disease in accordance with the invention herein disclosed, inter alia.

[0084] An agent that increases PP2A methylation and PP2A phosphatase activity is administered to a group of the hypercysteinemic mice. A placebo is administered to another group of the same mice. The group receiving the placebo exhibits significantly greater vascular inflamation and accelerated development of atherosclerosis, much as reported by Hofman et al. (March 2001).

The group receiving the agent does not exhibit significantly greater vascular inflamation and does not show the same accelerated development of atherosclerosis. The effect of the agent is seen in several experiments with different groups of mice and is dose dependent.

#### Example 15: Mouse in vivo PP2A methylation assays

[0085] Drug leads that test positive in the above screens are tested for efficacy in a mouse model for Alzheimer's. Briefly, mice are either fed a diet that contains the drug or the drug is administered by intra peritoneal injection, and effects on levels of PP2A methylation are assayed in brain using monoclonal antibodies that are specific for methylated PP2A (see above). In parallel with these studies we look for effects of altered levels of PP2A methylation on levels of phosphorylation of specific target proteins using monoclonal antibodies specific for epitopes that contain phosphorylation sites of interest. Drugs that have a beneficial effect on methylation and/or phosphorylation and on the subject mice are selected for additional studies.

# Example 16: Increased PP2A methylation decreases tau hyperphosphorylation in a mouse AD model

[0086] An agent that increases PP2A methylation and PP2A phosphatase activity is administered to groups of mice with hyperphosphorylated tau protein in a model of AD disease, including early onset. Other groups of the same mice are not treated, treated with placebo in the same way, or are treated with other agents that do not affect PP2A or related enzymes or factors. Each group of mice that is treated with the agent exhibits slowed increase of hyperphosphorylated tau, at least, and tau phosphorylation is seem to decrease in most of the mice, in some cases to normal levels. None of the mice receiving the agent before last stage AD, show further signs of AD development through the end of the experiment. Mice in the control groups, in contrast, show increasing tau phosphorylation and eventually develop the progressively more severe symptoms characteristic of these mouse AD models.

[0087] Hyperphosphorylation of MAP kinase proteins has been associated with neoplastic

transformation. We screen for agents that prevent elevated levels of MAP kinase

phosphorylation in the tissue culture and mouse screens (as described above). Agents that

prevent the elevated levels of MAP kinase phosphorylation that are induced by elevated total

plasma homocysteine are validated candidates for pharmaceuticals to treat or prevent cancer.

Example 18: Better levels of PPMS in genetically deficient mice

[0088] Mice genetically deficient in methylation metabolism and that, as a result, suffer from

deleterious PPMS, are kept on a normal diet. A placebo is administered to one group of the

mice. An agent that increases PP2A methylation and PP2A phosphatase activity is administered

to another group. Administration of the placebo and the agent is the same for each group except

for the presence of the agent in the composition administered to one group and not the other.

PPMS is determined for mice in each of the groups and the results are compared. Compared

with mice receiving the placebo, mice that receive the agent exhibit higher levels of PP2A

methylation, higher levels of PP2A phosphatase activity, and healthier levels of PPMS.

Example 19: Combinatorial library and HTS screen - Okadaic acid scaffolds

[0089] Okadaic acid is a potent inhibitor of both PP2A phosphatase activity and methylation of

PP2A. Moieties are identified in the compound that (1) most resemble structures likely to inhibit

the phosphatase activity of PP2A and (2) most resemble structures likely to inhibit PP2A

methylation. Putative methylation structures as described in (2) above, are evaluated and ranked

as to such considerations as the following: likelihood of inhibitory effect; likely strength thereof;

resemblance to other bioactive compounds and likelihood of unwanted effect; suitability as

scaffold for combinatorial synthesis; yield of likely inhibitory candidates if used as combinatorial

scaffold; ease of carrying out the syntheses necessary to obtain substantially the estimated yield;

and compound diversity yield space for each scaffold; among others.

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[0090] Based on the foregoing criteria, one or more Okadaic acid - derived scaffolds are

obtained and used for syntheses of a two-million compound library of derivatives to be screened

as methyltransferase and methylesterase inhibitors and agonists.

[0091] The compounds are screened in mixtures or individually in an HTS assay much as

described above. All mixtures showing positive results are sub-divided and retested. The

division and retesting is repeated for each sample giving a positive result, until as many of the

active candidates are singularly purified and then identified by MS (MALDI-TOF, for instance).

After retesting, a larger batch of each positive is prepared and then tested as described above for

activity in inhibiting or enhancing: PP2A phosphatase activity, methylation of PP2A, PP2A

MTase activity, and PP2A MEase activity. The candidates also are tested for deleterious effects

on cells in culture. Finally, the best candidates are tested in a mouse model of AD as described

above for all of the foregoing and for effect on tau hyperphosphorylation.

<u>REFERENCES</u>

The following references, keyed to the text numbers above, are herein incorporated by

reference in their entireties in parts pertinent to the aspects, embodiments, and practice of the

invention herein disclosed, particularly in parts pertinent to dietary and other disease models, to

PP2A and its activities, to methylation of PP2A, to assay and determination of phosphoprotein

phosphorylation, to drug discovery and assay relating to PP2A and other dephosphorylation

enzymes, and to methylation and demethylation enzymes active on the same, particularly those

that modify PP2A activities, to agents and their formulation regarding the same, and to

administration and treatment modalities affecting the same, as disclosed elsewhere herein,

among other things.

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